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#### (57) Abstract

The invention provides a nucleotide sequence representing a pathogenicity island found in species of pathogenic mycobacteria. The islands are shown as SEQ ID NOs: 3 and 4 and comprises several open reading frames encoding polypeptides. These polypeptides and their use in diagnosis and therapy form a further aspect of the invention.

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Novel polynucleotides and polypeptides in pathogenic mycobacteria and their use as diagnostics, vaccines and targets for chemotherapy.

This invention relates to the novel polynucleotide sequence we have designated "GS" which we have identified in pathogenic mycobacteria. GS is a pathogenicity island within 8kb of DNA comprising a core region of 5.75kb and an adjacent transmissable element within 2.25kb. GS is contained within Mycobacterium paratuberculosis, Mycobacterium avium subsp. silvaticum and some pathogenic isolates of M.avium. Functional portions of the core region of GS are also represented by regions with a high degree of homology that we have identified in cosmids containing genomic DNA from Mycobacterium tuberculosis.

## 15 Background to the invention

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Mycobacterium tuberculosis (Mtb) is a major cause of global diseases of humans as well as animals. Although conventional methods of diagnosis including microscopy, culture and skin testing exist for the recognition of these diseases, improved methods particularly new immunodiagnostics and detection systems are needed. Drugs used to treat tuberculosis are increasingly encountering the problem of resistant organisms. New drugs targeted at specific pathogenicity determinants as well as new vaccines for the prevention and treatment of tuberculosis are required. The importance of Mtb as a global pathogen is reflected in the commitment being made to sequencing the entire genome of this organism. This has generated a large amount of DNA sequence data of genomic DNA within cosmid and other libraries. Although the DNA sequence is known in the art, the functions of the vast majority of these sequences, the proteins they encode, the biological significance of these proteins, and the overall relevance and use of these genes and their products as diagnostics, vaccines and targets for chemotherapy for tuberculous disease, remains entirely unknown.

35 Mycobacterium avium subsp.silvaticum (Mavs) is a pathogenic mycobacterium causing diseases of animals and birds, but it can

also affect humans. Mycobacterium paratuberculosis (Mptb) causes chronic inflammation of the intestine in many species of animals including primates and can also cause Crohn's disease in humans. Mptb is associated with other chronic inflammatory diseases of Subclinical Mptb infection is humans such as sarcoidosis. widespread in domestic livestock and is present in milk from organism is more resistant infected animals. The pasteurisation than Mtb and can be conveyed to humans in retail Mptb is also present in water supplies, milk supplies. particularly those contaminated with run-off from heavily grazed pastures. Mptb and Mavs contain the insertion elements IS900 and IS902 respectively, and these are linked to pathogenicity in IS900 and IS902 provide convenient highly these organisms. specific multi-copy DNA targets for the sensitive detection of these organisms using DNA-based methods and for the diagnosis of infections in animals and humans. Much improvement is however required in the immunodiagnosis of Mptb and Mavs infections in animals and humans. Mptb and Mavs are in general, resistant in vivo to standard anti-tuberculous drugs. Although substantial clinical improvements in infections caused by Mptb, such as may result from treatment of patients with Crohn's disease, combinations of existing drugs such as Rifabutin, Clarithromycin additional effective drug treatments are or Azithromycin, Furthermore, there is an urgent need for effective required. vaccines for the prevention and treatment of Mptb and Mavs infections in animals and humans based upon the recognition of specific pathogenicity determinants.

Pathogenicity islands are, in general, 7-9kb regions of DNA comprising a core domain with multiple ORFs and an adjacent transmissable element. The transmissable element also encodes proteins which may be linked to pathogenicity, such as by providing receptors for cellular recognition. Pathogenicity islands are envisaged as mobile packages of DNA which, when they enter an organism, assist in bringing about its convertion from a non-disease-causing to a disease-causing strain.

## Description of the Drawings

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Figure 1(a) and (b) shows a linear map of the pathogenicity island GS in Mavs (Fig 1a) and in Mptb (Fig 1b). The main open reading frames are illustrated as ORFs A to H. ORFs A to F are found within the core region of GS. ORFs G and H are encoded by the adjacent transmissable element portion of GS.

## Disclosure of the invention

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Using a DNA-based differential analysis technology we have discovered and characterised a novel polynucleotide in Mptb (isolates 0022 from a Guernsey cow and 0021 from a red deer). This polynucleotide comprises the gene region we have designated GS is found in Mptb using the identifier DNA sequences Seq.ID.No 1 and 2 where the Seq.ID No2 is the complementary sequence of Seq.ID No 1. GS is also identified in Mavs. complete DNA sequence incorporating the positive strand of GS from an isolate of Mavs comprising 7995 nucleotides, including the core region of GS and adjacent transsmissable element, DNA sequence comprising 4435 bp of the given in Seq.ID No.3. positive strand of GS obtained from an isolate of Mptb including the core region of GS (nucleotides 1614 to 6047 of GS in Mavs) is given in Seq.ID No 4. The DNA sequence of GS from Mptb is highly (99.4%) homologous to GS in Mavs. The remaining portion of the DNA sequence of GS in Mptb, is readily obtainable by a person skilled in the art using standard laboratory procedures. The entire functional DNA sequence including core region and transmisable element of GS in Mptb and Mavs as described above, comprise the polynucleotide sequences of the invention.

There are 8 open reading frames (ORFs) in GS. Six of these designated GSA, GSB, GSC, GSD, GSE and GSF are encoded by the core DNA region of GS which, characteristically for a pathogenicity island, has a different GC content than the rest of the microbial genome. Two ORFs designated GSG and GSH are encoded by the transmissable element of GS whose GC content resembles that of the rest of the mycobacterial genome. The ORF GSH comprises two sub-ORFs  $\rm H_1$   $\rm H_2$  on the complementary DNA strand linked by a programmed frameshifting site so that a single polypeptide is translated from the ORF GSH. The nucleotide

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sequences of the 8 ORFs in GS and their translations are shown in Seq. ID No 5 to Seq.ID No 29 as follows:

- ORF A: Seq. ID No 5 Nucleotides 50 to 427 of GS from Mavs Seq. ID No 6 Amino acid sequence encoded by Seq. ID No 5.
  - ORF B: Seq. ID No 7 Nucleotides 772 to 1605 of GS from Mavs Seq. ID No 8 Amino acid sequence encoded by Seq. ID No 7.
- ORF C: Seq. ID No 9 Nucleotides 1814 to 2845 of GS from Mavs

  Seq. ID No 10 Amino acid sequence encoded by Seq. ID No

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  Seq. ID No 11 Nucleotides 201 to 1232 of GS from Mptb

  Seq. ID No 12 Amino acid sequence encoded by Seq. ID No
- 15 ORF D: Seq. ID No 13 Nucleotides 2785 to 3804 of GS from Mavs Seq. ID No 14 Amino acid sequence encoded by Seq.ID No 13.

  Seq. ID No 15 Nucleotides 1172 to 2191 of GS from Mpth Seq. ID No 16 Amino acid sequence encoded by Seq.ID No 15.
  - ORF E: Seq. ID No 17 Nucleotides 4080 to 4802 of GS from Mavs Seq. ID No 18 Amino acid sequence encoded by Seq.ID No 17.

    Seq. ID No 19 Nucleotides 2467 to 3189 of GS from Mptb Seq. ID No 20 Amino acid sequence encoded by Seq.ID No
  - ORF F: Seq. ID No 21 Nucleotides 4947 to 5747 of GS from Mavs Seq. ID No 22 Amino acid sequence encoded by Seq. ID No 21.
- Seq. ID No 23 Nucleotides 3335 to 4135 of GS from Mptb Seq. ID No 24 Amino acid sequence encoded by Seq. ID No 23.

- ORF G: Seq. ID No 25 Nucleotides 6176 to 7042 of GS from Mavs Seq. ID No 26 Amino acid sequence encoded by Seq.ID No 25.
- ORF H: Seq.ID No 27 Nucleotides 7953 to 6215 from Mavs.
- 5 ORF  $H_1$ : Seq.ID No 28 Amino acid sequence encoded by nucleotides 7953 to 7006 of Seq.ID No 27
  - ORF  $H_2$ : Seq.ID No 29 Amino acid sequence encoded by nucleotides 7009 to 6215 of Seq.ID No 27
- The polynucleotides in Mtb with homology to the ORFs B, C, E and 10 F of GS in Mptb and Mavs, and the polypeptides they are now known to encode as a result of our invention, are as follows:

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- ORF B: Seq.ID No 30 Cosmid MTCY277 nucleotides 35493 to 34705 Seq.ID No 31 Amino acid sequence encoded by Seq.ID No30.
- ORF C: Seq.ID No 32 Cosmid MTCY277 nucleotides 31972 to 32994 Seq.ID No 33 Amino acid sequence encoded by Seq.ID No32.
- ORF E: Seq.ID No 34 Cosmid MTCY277 nucleotides 34687 to 33956

  Seq.ID No 35 Amino acid sequence encoded by Seq.ID No34.
  - ORF E: Seq.ID No 36 Cosmid MTO24 nucleotides 15934 to 15203 Seq.ID No 37 Amino acid sequence encoded by Seq.ID No36.
- 25 ORF F: Seq.ID No38 Cosmid MTO24 nucleotides 15133 to 14306 Seq.ID No 39 Amino acid sequence encoded by Seq.ID No38.

The proteins and peptides encoded by the ORFs A to H in Mptb and Mavs and the amino acid sequences from homologous genes we have

discovered in Mtb given in Seq.ID Nos 31, 33, 35, 37 and 39, as described above and fragments thereof, comprise the polypeptides of the invention. The polypeptides of the invention are believed to be associated with specific immunoreactivity and with the pathogenicity of the host micro-organisms from which they were obtained.

The present invention thus provides a polynucleotide in substantially isolated form which is capable of selectively hybridising to sequence ID Nos 3 or 4 or a fragment thereof. The polynucleotide fragment may alternatively comprise a sequence selected from the group of Seq.ID.No: 5, 7, 9, 11, 13, 15, 17, 19, 21, 23, 25 and 27. The invention further provides a polynucleotide in substantially isolated form whose sequence consists essentially of a sequence selected from the group Seq ID Nos. 30, 32, 34, 36 and 38, or a corresponding sequence selectively hybridizable thereto, or a fragment of said sequence or corresponding sequence.

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The invention further provides diagnostic probes such as a probe which comprises a fragment of at least 15 nucleotides of a polynucleotide of the invention, or a peptide nucleic acid or similar synthetic sequence specific ligand, optionally carrying a revealing label. The invention also provides a vector carrying a polynucleotide as defined above, particularly an expression vector.

The invention further provides a polypeptide in substantially 25 isolated form which comprises any one of the sequences selected from the group consisting Seq. ID. No: 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 29, 31, 33, 35, 37 and 39, or a polypeptide substantially homologous thereto. The invention additionally provides a polypeptide fragment which comprises a fragment of a 30 polypeptide defined above, said fragment comprising at least 10 The invention also provides amino acids and an epitope. polynucleotides in substantially isolated form which encode polypeptides of the invention, and vectors which comprise such polynucleotides, as well as antibodies capable of binding such 35 polypeptides. In an additional aspect, the invention provides

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kits comprising polynucleotides, polypeptides, antibodies or synthetic ligands of the invention and methods of using such kits in diagnosing the presence or absence of mycobacteria in a sample. The invention also provides pharmaceutical compositions comprising polynucleotides of the invention, polypeptides of the invention or antisense probes and the use of such compositions prevention of treatment ordiseases caused The invention also provides polynucleotihe mycobacteria. prevention and treatment of infections due to GS-containing pathogenic mycobacteria in animals and humans and as a means of in vivo susceptibility of said mycobacteria The invention also provides bacteria or antimicrobial drugs. viruses transformed with polynucleotides of the invention for use The invention further provides Mptb or Mavs as vaccines. which all or part or the polynucleotides of the invention have been deleted or disabled to provide mutated organisms of lower pathogenicity for use as vaccines in animals and humans. invention further provides Mtb in which all or part of the polynucleotides encoding polypeptides of the invention have been deleted or disabled to provided mutated organisms or lower pathogenicity for use as vaccines in animals and humans.

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A further aspect of the invention is our discovery of homologies between the ORFs B. C and E in GS on the one hand, and Mtb cosmid MTCY277 on the other (data from Genbank database using the computer programmes BLAST and BLIXEM). The homologous ORFs in MTCY277 are adjacent to one another consistent with the form of another pathogenicity island in Mtb. A further aspect of the invention is our discovery of homologies between ORFs E and F in GS, and Mtb cosmid MTO24 (also Genbank, as above) with the homologous ORFs close to one another. The use of polynucleotides and polypeptides from Mtb (Seq. ID Nos 30,31, 32, 33, 34, 35, 36, 37, 38 and 39) in substantially isolated form as diagnostics, vaccines and targets for chemotherapy, for the management and prevention of Mtb infections in humans and animals, and the involved in the preparation and use of processes diagnostics, vaccines and new chemotherapeutic agents, comprise further aspects of the invention.

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## Detailed description of the invention.

## A. Polynucleotides

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Polynucleotides of the invention as defined herein may comprise DNA or RNA. They may also be polynucleotides which include within them synthetic or modified nucleotides or peptide nucleic A number of different types of modification to oligonucleotides are known in the art. These methylphosphonate and phosphorothioate backbones, addition of acridine or polylysine chains at the 3' and/or 5' ends of the molecule. For the purposes of the present invention, it is to be understood that the polynucleotides described herein may be modified by any method available in the art. Such modifications may be carried out in order to couple the said polynucleotide to a solid phase or to enhance the recognition, the in vivo activity, or the lifespan of polynucleotides of the invention.

A number of different types of polynucleotides of the invention are envisaged. In the broadest aspect, polynucleotides and fragments thereof capable of hybridizing to SEQ ID NO:3 or 4 form a first aspect of the invention. This includes the polynucleotide of SEQ ID NO: 3 or 4. Within this class of polynucleotides various sub-classes of polynucleotides are of particular interest.

One sub-class of polynucleotides which is of interest is the class of polynucleotides encoding the open reading frames A, B, C, D, E, F, G and H, including SEQ ID NOs:5, 7, 9, 11, 13, 15, 17, 19, 21, 23, 25 and 27. As discussed below, polynucleotides encoding ORF H include the polynucleotide sequences 7953 to 7006 and 7009 to 6215 within SEQ ID NO: 27, as well as modified sequences in which the frame-shift has been modified so that the two sub-reading frames are placed in a single reading frame. This may be desirable where the polypeptide is to be produced in recombinant expression systems.

The invention thus provides a polynucleotide in substantially isolated form which encodes any one of these ORFs or combinations

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thereof. Combinations thereof includes combinations of 2, 3, 4, 5 or all of the ORFs. Polynucleotides may be provided which comprise an individual ORF carried in a recombinant vector including the vectors described herein. Thus in one preferred aspect the invention provides a polynucleotide in substantially isolated form capable of selectively hybridizing to the nucleic acid comprising ORFs A to F of the core region of the Mptb and Mavs pathogenicity islands of the invention. Fragments thereof corresponding to ORFs A to E, B to F, A to D, B to E, A to C, B to D or any two adjacent ORFs are also included in the invention.

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Polynucleotides of the invention will be capable of selectively hybridizing to the corresponding portion of the GS region, or to the corresponding ORFs of Mtb described herein. "selectively hybridizing" indicates that the polynucleotides will hybridize, under conditions of medium to high stringency (for example 0.03 M sodium chloride and 0.03 M sodium citrate at from about 50°C to about 60°C) to the corresponding portion of SEQ ID NO:3 or 4 or the complementary strands thereof but not to genomic DNA from mycobacteria which are usually non-pathogenic including non-pathogenic species of M.avium. Such polynucleotides will generally be generally at least 68%, e.g. at least 70%, preferably at least 80 or 90% and more preferably at least 95% homologous to the corresponding DNA of GS. The corresponding portion will be of over a region of at least 20, preferably at least 30, for instance at least 40, 60 or 100 or more contiguous nucleotides.

By "corresponding portion" it is meant a sequence from the GS region of the same or substantially similar size which has been determined, for example by computer alignment, to have the greatest degree of homology to the polynucleotide.

Any combination of the above mentioned degrees of homology and minimum sizes may be used to define polynucleotides of the invention, with the more stringent combinations (i.e. higher homology over longer lengths) being preferred. Thus for example a polynucleotide which is at least 80% homologous over 25, preferably 30 nucleotides forms one aspect of the invention, as

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does a polynucleotide which is at least 90% homologous over 40 nucleotides.

A further class of polynucleotides of the invention is the class of polynucleotides encoding polypeptides of the invention, the 5 polypeptides of the invention being defined in section B below. the redundancy of the genetic code polynucleotides may be of a lower degree of homology than required for selective hybridization to the GS region. when such polynucleotides encode polypeptides of the invention 10 these polynucleotides form a further aspect. It may for example be desirable where polypeptides of the invention are produced recombinantly to increase the GC content of such polynucleotides. This increase in GC content may result in higher levels of expression via codon usage more appropriate to the host cell in which recombinant expression is taking place.

An additional class of polynucleotides of the invention are those obtainable from cosmids MTCY277 and MT024 (containing Mtb genomic sequences), which polynucleotides consist essentially of the fragment of the cosmid containing an open reading frame encoding any one of the homologous ORFs B, C, E or F respectively. Such polynucleotides are referred to below as Mtb polynucleotides. However, where reference is made to polynucleotides in general such reference includes Mtb polynucleotides unless the context is explicitly to the contrary. In addition, the invention provides polynucleotides which encode the same polypeptide as the abovementioned ORFs of Mtb but which, due to the redundancy of the genetic code, have different nucleotide sequences. These form further Mtb polynucleotides of the invention. Fragments of Mtb polynucleotides suitable for use as probes or primers also form a further aspect of the invention.

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The invention further provides polynucleotides in substantially isolated form capable of selectively hybridizing (where selectively hybridizing is as defined above) to the Mtb polynucleotides of the invention.

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The invention further provides the Mtb polynucleotides of the invention linked, at either the 5' and/or 3' end to polynucleotide sequences to which they are not naturally contiguous. Such sequences will typically be sequences found in cloning or expression vectors, such as promoters, 5' untranslated sequence, 3' untranslated sequence or termination sequences. The sequences may also include further coding sequences such as signal sequences used in recombinant production of proteins.

Further polynucleotides of the invention are illustrated in the accompanying examples.

Polynucleotides of the invention may be used to produce a primer, e.g. a PCR primer, a primer for an alternative amplification reaction, a probe e.g. labelled with a revealing label by conventional means using radioactive or non-radioactive labels or a probe linked covalently to a solid phase, or the polynucleotides may be cloned into vectors. Such primers, probes and other fragments will be at least 15, preferably at least 20, for example at least 25, 30 or 40 or more nucleotides in length, and are also encompassed by the term polynucleotides of the invention as used herein.

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Primers of the invention which are preferred include primers directed to any part of the ORFs defined herein. The ORFs from other isolates of pathogenic mycobacteria which contain a GS region may be determined and conserved regions within each individual ORF may be identified. Primers directed to such conserved regions form a further preferred aspect of the invention. In addition, the primers and other polynucleotides of the invention may be used to identify, obtain and isolate ORFs capable of selectively hybridizing to the polynucleotides of the invention which are present in pathogenic mycobacteria but which are not part of a pathogenicity island in that particular species of bacteria. Thus in addition to the ORFs B, C, E and F which have been identified in Mtb, similar ORFs may be identified in other pathogens and ORFs corresponding to the GS ORFs C, D, E, F and H, may also be identified.

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Polynucleotides such as DNA polynucleotides and probes according to the invention may be produced recombinantly, synthetically, or by any means available to those of skill in the art. They may also be cloned by standard techniques.

In general, primers will be produced by synthetic means, involving a step-wise manufacture of the desired nucleic acid sequence one nucleotide at a time. Techniques for accomplishing this using automated techniques are readily available in the art. Longer polynucleotides will generally be produced 10 recombinant means, for example using a PCR (polymerase chain reaction) cloning techniques. This will involve making a pair or primers (e.g. of about 15-30 nucleotides) to a region of GS, which it is desired to clone, bringing the primers into contact with genomic DNA from a mycobacterium or a vector carrying the 15 GS sequence, performing a polymerase chain reaction under conditions which bring about amplification of the desired region, isolating the amplified fragment (e.g. by purifying the reaction mixture on an agarose gel) and recovering the amplified DNA. primers may be designed to contain suitable restriction enzyme recognition sites so that the amplified DNA can be cloned into 20 a suitable cloning vector.

Such techniques may be used to obtain all or part of the GS or ORF sequences described herein, as well as further genomic clones containing full open reading frames. Although in general such techniques are well known in the art, reference may be made in particular to Sambrook J., Fritsch EF., Maniatis T (1989). Molecular cloning: a Laboratory Manual, 2nd edn. Cold Spring Harbor, New York, Cold Spring Harbor Laboratory.

Polynucleotides which are not 100% homologous to the sequences 30 of the present invention but fall within the scope of the invention can be obtained in a number of ways.

Other isolates or strains of pathogenic mycobacteria will be expected to contain allelic variants of the GS sequences described herein, and these may be obtained for example by probing genomic DNA libraries made from such isolates or strains

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of bacteria using GS or ORF sequences as probes under conditions of medium to high stringency (for example 0.03M sodium chloride and 0.03M sodium citrate at from about 50°C to about 60°C).

A particularly preferred group of pathogenic mycobacteria are isolates of *M.paratuberculosis*. Polynucleotides based on GS regions from such bacteria are particularly preferred. Preferred fragments of such regions include fragments encoding individual open reading frames including the preferred groups and combinations of open reading frames discussed above.

Alternatively, such polynucleotides may be obtained by site 10 directed mutagenesis of the GS or ORF sequences or allelic variants thereof. This may be useful where for example silent codon changes are required to sequences to optimise codon cell ìn which particular host preferences for a polynucleotide sequences are being expressed. Other sequence 15 changes may be desired in order to introduce restriction enzyme recognition sites, or to alter the property or function of the polypeptides encoded by the polynucleotides of the invention. Such altered property or function will include the addition of amino acid sequences of consensus signal peptides known in the 20 art to effect transport and secretion of the modified polypeptide Another altered property will include of the invention. metagenesis of a catalytic residue or generation of fusion proteins with another polypeptide. Such fusion proteins may be 25 with an enzyme, with an antibody or with a cytokine or other ligand for a receptor, to target a polypeptide of the invention to a specific cell type in vitro or in vivo.

The invention further provides double stranded polynucleotides comprising a polynucleotide of the invention and its complement.

30 Polynucleotides or primers of the invention may carry a revealing label. Suitable labels include radioisotopes such as <sup>32</sup>P or <sup>35</sup>S, enzyme labels, other protein labels or smaller labels such as biotin or fluorophores. Such labels may be added to polynucleotides or primers of the invention and may be detected using by techniques known per se.

Polynucleotides or primers of the invention or fragments thereof labelled or unlabelled may be used by a person skilled in the art in nucleic acid-based tests for the presence or absence of Mptb, Mavs, other GS-containing pathogenic mycobacteria, or Mtb applied to samples of body fluids, tissues, or excreta from animals and humans, as well as to food and environmental samples such as river or ground water and domestic water supplies.

Human and animal body fluids include sputum, blood, serum, plasma, saliva, milk, urine, csf, semen, faeces and infected discharges. Tissues include intestine, mouth ulcers, skin, lymph nodes, spleen, lung and liver obtained surgically or by a biopsy technique. Animals particularly include commercial livestock such as cattle, sheep, goats, deer, rabbits but wild animals and animals in zoos may also be tested.

Such tests comprise bringing a human or animal body fluid or 15 tissue extract, or an extract of an environmental or food sample, into contact with a probe comprising a polynucleotide or primer of the invention under hybridising conditions and detecting any duplex formed between the probe and nucleic acid in the sample. Such detection may be achieved using techniques such as PCR or 20 by immobilising the probe on a solid support, removing nucleic acid in the sample which is not hybridized to the probe, and then detecting nucleic acid which has hybridized to the probe. Alternatively, the sample nucleic acid may be immobilized on a solid support, and the amount of probe bound to such a support 25 Suitable assay methods of this any other can be detected. formats can be found in for example WO89/03891 and WO90/13667.

Polynucleotides of the invention or fragments thereof labelled or unlabelled may also be used to identify and characterise different strains of Mptb, Mavs, other GS-containing pathogenic mycobacteria, or Mtb, and properties such as drug resistance or susceptibility.

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The probes of the invention may conveniently be packaged in the form of a test kit in a suitable container. In such kits the probe may be bound to a solid support where the assay format for

which the kit is designed requires such binding. The kit may also contain suitable reagents for treating the sample to be probed, hybridising the probe to nucleic acid in the sample, control reagents, instructions, and the like.

The use of polynucleotides of the invention in the diagnosis of inflammatory diseases such as Crohn's disease or sarcoidosis in humans or Johne's disease in animals form a preferred aspect of the invention. The polynucleotides may also be used in the prognosis of these diseases. For example, the response of a human or animal subject in response to antibiotic, vaccination or other therapies may be monitored by utilizing the diagnostic methods of the invention over the course of a period of treatment and following such treatment.

The use of Mtb polynucleotides (particularly in the form of probes and primers) of the invention in the above-described methods form a further aspect of the invention, particularly for the detection, diagnosis or prognosis of Mtb infections.

#### B. Polypeptides.

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the invention include polypeptides Polypeptides of substantially isolated form encoded by GS. This includes the length polypeptides encoded by the positive full complementary negative strands of GS. Each of the full length polypeptides will contain one of the amino acid sequences set out in Seq ID NOs:6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28 and Polypeptides of the invention further include variants of such sequences, including naturally occurring allelic variants and synthetic variants which are substantially homologous to said polypeptides. In this context, substantial homology is regarded as a sequence which has at least 70%, e.g. 80%, 90%, 95% or 98% amino acid homology (identity) over 30 or more, e.g 40, 50 or 100 amino acids. For example, one group of substantially homolgous polypeptides are those which have at least 95% amino acid identity to a polypeptide of any one of Seq ID NOs:6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28 and 29 over their entire length. Even more preferably, this homology is 98%.

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Polypeptides of the invention further include the polypeptide sequences of the homologous ORFs of Mtb, namely Seq ID Nos. 31, 33, 35, 37 and 39. Unless explicitly specified to the contrary, reference to polypeptides of the invention and their fragments include these Mtb polypeptides and fragments, and variants thereof (substanially homologous to said sequences) as defined herein.

Polypeptides of the invention may be obtained by the standard techniques mentioned above. Polypeptides of the invention also include fragments of the above mentioned full length polypeptides and variants thereof, including fragments of the sequences set out in SEQ ID NOs:6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 29, 31, 33, 35, 37 and 39. Such fragments for example of 8, 10, 12, 15 or up to 30 or 40 amino acids may also be obtained synthetically using standard techniques known in the art.

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Preferred fragments include those which include an epitope, especially an epitope which is specific to the pathogenicity of the mycobacterial cell from which the polypeptide is derived. Suitable fragments will be at least about 5, e.g. 8, 10, 12, 15 20 or 20 amino acids in size, or larger. Epitopes may be determined either by techniques such as peptide scanning techniques as described by Geysen et al, Mol.Immunol., 23; 709-715 (1986), as well as other techniques known in the art.

The term "an epitope which is specific to the pathogenicity of the mycobacterial cell" means that the epitope is encoded by a portion of the GS region, or by the corresponding ORF sequences of Mtb which can be used to distinguish mycobacteria which are pathogenic by from related non-pathogenic mycobacteria including non-pathogenic species of M. avium. This may be determined using routine methodology. A candidate epitope from an ORF may be prepared and used to immunise an animal such as a rat or rabbit in order to generate antibodies. The antibodies may then be used to detect the presence of the epitope in pathogenic mycobacteria and to confirm that non-pathogenic mycobacteria do not contain any proteins which react with the epitope. Epitopes may be 35 linear or conformational.

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Polypeptides of the invention may be in a substantially isolated form. It will be understood that the polypeptide may be mixed with carriers or diluents which will not interfere with the intended purpose of the polypeptide and still be regarded as substantially isolated. A polypeptide of the invention may also be in a substantially purified form, in which case it will generally comprise the polypeptide in a preparation in which more than 90%, e.g. 95%, 98% or 99% of the polypeptide in the preparation is a polypeptide of the invention.

10 Polypeptides of the invention may be modified to confer a desired property or function for example by the addition of Histidine residues to assist their purification or by the addition of a signal sequence to promote their secretion from a cell.

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Thus, polypeptides of the invention include fusion proteins which comprise a polypeptide encoding all or part of one or more of an ORF of the invention fused at the N- or C-terminus to a second sequence to provide the desired property or function. Sequences which promote secretion from a cell include, for example the yeast  $\alpha$ -factor signal sequence.

A polypeptide of the invention may be labelled with a revealing label. The revealing label may be any suitable label which allows the polypeptide to be detected. Suitable labels include radioisotopes, e.g. <sup>125</sup>I, <sup>35</sup>S enzymes, antibodies, polynucleotides and ligands such as biotin. Labelled polypeptides of the invention may be used in diagnostic procedures such as immunoassays in order to determine the amount of a polypeptide of the invention in a sample. Polypeptides or labelled polypeptides of the invention may also be used in serological or cell mediated immune assays for the detection of immune reactivity to said polypeptides in animals and humans using standard protocols.

A polypeptide or labelled polypeptide of the invention or fragment thereof may also be fixed to a solid phase, for example the surface of an immunoassay well, microparticle, dipstick or biosensor. Such labelled and/or immobilized polypeptides may be

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packaged into kits in a suitable container along with suitable reagents, controls, instructions and the like.

Such polypeptides and kits may be used in methods of detection of antibodies or cell mediated immunoreactivity, to 5 mycobacterial proteins and peptides encoded by the ORFs of the invention and their allelic variants and fragments, using Such host antibodies or cell mediated immune immunoassay. reactivity will occur in humans or animals with an immune system which detects and reacts against polypeptides of the invention. The antibodies may be present in a biological sample from such humans or animals, where the biological sample may be a sample as defined above particularly blood, milk or saliva.

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Immunoassay methods are well known in the art and will generally comprise:

- providing a polypeptide of the invention comprising an (a) epitope bindable by an antibody against said mycobacterial polypeptide;
- incubating a biological sample with said polypeptide (b) under conditions which allow for the formation of an antibody-antigen complex; and
- whether antibody-antigen complex (c) determining comprising said polypeptide is formed.

Immunoassay methods for cell mediated immune reactivity in animals and humans are also well known in the art (e.g. as described by Weir et al 1994, J. Immunol Methods 176; 93-101) and will generally comprise

- providing a polypeptide of the invention comprising an epitope bindable by a lymphocyte or macrophage or other cell receptor;
- incubating a cell sample with said polypeptide under (b) conditions which allow for a cellular immune response such as release of cytokines or other mediator to occur; and
- detecting the presence of said cytokine or mediator in the incubate.

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Polypeptides of the invention may be made by standard synthetic means well known in the art or recombinantly, as described below.

Polypeptides of the invention or fragments thereof labelled or unlabelled may also be used to identify and characterise different strains of Mptb, Mavs, other GS-containing pathogenic mycobacteria, or Mtb, and properties such as drug resistance or susceptibility.

The polypeptides of the invention may conveniently be packaged in the form of a test kit in a suitable container. In such kits the polypeptide may be bound to a solid support where the assay format for which the kit is designed requires such binding. The kit may also contain suitable reagents for treating the sample to be examined, control reagents, instructions, and the like.

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The use of polypeptides of the invention in the diagnosis of inflammatory diseases such as Crohn's disease or sarcoidosis in humans or Johne's disease in animals form a preferred aspect of the invention. The polypeptides may also be used in the prognosis of these diseases. For example, the response of a human or animal subject in response to antibiotic or other therapies may be monitored by utilizing the diagnostic methods of the invention over the course of a period of treatment and following such treatment.

The use of *Mtb* polypeptides of the invention in the above-described methods form a further aspect of the invention, particularly for the detection, diagnosis or prognosis of *Mtb* infections.

Polypeptides of the invention may also be used in assay methods for identifying candidate chemical compounds which will be useful in inhibiting, binding to or disrupting the function of said polypeptides required for pathogenicity. In general, such assays involve bringing the polypeptide into contact with a candidate inhibitor compound and observing the ability of the compound to disrupt, bind to or interfer with the polypeptide.

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There are a number of ways in which the assay may be formatted. For example, those polypeptides which have an enzymatic function may be assayed using labelled substrates for the enzyme, and the amount of, or rate of, conversion of the substrate into a product 5 measured, e.g by chromatograpy such as HPLC or by a colourimetric assay. Suitable labels include 35S, 125I, biotin or enzymes such as horse radish peroxidase.

For example, the gene product of ORF C is believed to have GDPmannose dehydratase activty. Thus an assay for inhbitors of the gene product may utilise for example labelled GDP-mannose, GDP or mannose and the activity of the gene product followed. D encodes a gene related to the synthesis and regulation of capuslar polysaccharides, which are often associated with pathogenicity. Labelled polysaccharide invasiveness and substrates may be used in assays of the ORF D gene product. gene product of ORF F encodes a protein with putative glucosyl transferase activity and thus labelled amino sugars such as  $\beta$ -1-3-N-acetylglucosamine may be used as substrates in assays.

Candidate chemical compounds which may be used may be natural or 20 synthetic chemical compounds used in drug screening programmes. Extracts of plants which contain several characterised or uncharacterised components may also be used.

Alternatively, the a polypeptide of the invention may be screened against a panel of peptides, nucleic acids or other chemical functionalities which are generated by combinatorial chemistry. This will allow the definition of chemical entities which bind to polypeptides of the invention. Typically, the polypeptide of the invention will be brought into contact with a panel of compounds from a combinantorial library, with either the panel or the polypeptide being immobilized on a solid phase, under conditions suitable for the polypeptide to bind to the panel. The solid phase will then be washed under conditions in which only specific interactions between the polypeptide and individual members of the panel are retained, and those specific members may 35 be utilized in further assays or used to design further panels of candidate compounds.

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For example, a number of assay methods to define peptide interaction with peptides are known. For example, WO86/00991 describes a method for determining mimotopes which comprises making panels of catamer preparations, for example octamers of amino acids, at which one or more of the positions is defined and the remaining positions are randomly made up of other amino acids, determining which catamer binds to a protein of interest and re-screening the protein of interest against a further panel based on the most reactive catamer in which one or more additional designated positions are systematically varied. This may be repeated throughout a number of cycles and used to build up a sequence of a binding candidate compound of interest.

WO89/03430 describes screening methods which permit the preparation of specific mimotopes which mimic the immunological activity of a desired analyte. These mimotopes are identified by reacting a panel of individual peptides wherein said peptides are of systematically varying hydrophobicity, amphipathic characteristics and charge patterns, using an antibody against an antigen of interest. Thus in the present case antibodies against the a polypeptide of the inventoin may be employed and mimotope peptides from such panels may be identified.

#### C. Vectors.

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Polynucleotides of the invention can be incorporated into a recombinant replicable vector. The vector may be used to replicate the nucleic acid in a compatible host cell. Thus in a further embodiment, the invention provides a method of making polynucleotides of the invention by introducing a polynucleotide of the invention into a replicable vector, introducing the vector into a compatible host cell, and growing the host cell under conditions which bring about replication of the vector. The vector may be recovered from the host cell. Suitable host cells are described below in connection with expression vectors.

## D. Expression Vectors.

Preferably, a polynucleotide of the invention in a vector is operably linked to a control sequence which is capable of providing for the expression of the coding sequence by the host cell, i.e. the vector is an expression vector. The term "operably linked" refers to a juxtaposition wherein the components described are in a relationship permitting them to function in their intended manner. A control sequence "operably linked" to a coding sequence is ligated in such a way that expression of the coding sequence is achieved under conditions compatible with the control sequences. Such vectors may be transformed into a suitable host cell as described above to provide for expression of a polypeptide of the invention. Thus, in a further aspect the invention provides a process for preparing polypeptides according to the invention which comprises cultivating a host cell transformed or transfected with an expression vector as described above, under conditions to provide for expression by the vector of a coding sequence encoding the polypeptides, and recovering the expressed polypeptides.

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A further embodiment of the invention provides vectors for the replication and expression of polynucleotides of the invention, or fragments thereof. The vectors may be for example, plasmid, virus or phage vectors provided with an origin of replication, optionally a promoter for the expression of the polynucleotide and optionally a regulator of the promoter. vectors may contain one or more selectable marker genes, for example an ampicillin resistance gene in the case of a bacterial plasmid or a neomycin resistance gene for a mammalian vector. Vectors may be used in vitro, for example for the production of RNA or used to transfect or transform a host cell. The vector may also be adapted to be used in vivo, for example in a method of naked DNA vaccination or gene therapy. A further embodiment of the invention provides host cells transformed or transfected with the vectors for the replication and expression of polynucleotides of the invention, including the DNA of GS, the open reading frames thereof and other corresponding ORFs particularly ORFs B, C, E and F from Mtb. The cells will be chosen to be compatible with the said vector and may for example be bacterial, yeast, insect or mammalian.

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Expression vectors are widely available in the art and can be obtained commercially. Mammalian expression vectors may comprise a mammalian or viral promoter. Mammalian promoters include the metallothionien promoter. Viral promoters include promoters from adenovirus, the SV40 large T promoter and retroviral LTR promoters. Promoters compatible with insect cells include the polyhedrin promoter. Yeast promoters include the alcohol dehydrogenase promoter. Bacterial promoters include the  $\beta$ -galactosidase promoter.

10 The expression vectors may also comprise enhancers, and in the case of eukaryotic vectors polyadenylation signal sequence downstream of the coding sequence being expressed.

Polypeptides of the invention may be expressed in suitable host cells, for example bacterial, yeast, plant, insect and mammalian cells, and recovered using standard purification techniques including, for example affinity chromatography, HPLC or other chromatographic separation techniques.

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Polynucleotides according to the invention may also be inserted into the vectors described above in an antisense orientation in order to provide for the production of antisense RNA. Antisense RNA or other antisense polynucleotides or ligands may also be produced by synthetic means. Such antisense polynucleotides may be used in a method of controlling the levels of the proteins encoded by the ORFs of the invention in a mycobacterial cell.

25 Polynucleotides of the invention may also be carried by vectors suitable for gene therapy methods. Such gene therapy methods include those designed to provide vaccination against diseases caused by pathogenic mycobacteria or to boost the immune response of a human or animal infected with a pathogenic mycobacteria.

30 For example, Ziegner et al, AIDS, 1995, 9;43-50 describes the use of a replication defective recombinant amphotropic retrovirus to boost the immune response in patients with HIV infection. Such a retrovirus may be modified to carry a polynucleotide encoding a polypeptide or fragment thereof of the invention and the

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retrovirus delivered to the cells of a human or animal subject in order to provide an immune response against said polypeptide. The retrovirus may be delivered directly to the patient or may be used to infecte cells ex-vivo, e.g. fibroblast cells, which are then introduced into the patient, optionally after being inactivated. The cells are desirably autologous or HLA-matched cells from the human or animal subject.

Gene therapy methods including methods for boosting an immune response to a particluar pathogen are disclosed generally in for example WO95/14091, the disclosure of which is incoporated herein Recombinant viral vectors include retroviral by reference. vectors, adenoviral vectors, adeno-associated viral vectors, vaccinia virus vectors, herpes virus vectors and alphavirus Alpha virus vectors are described in, for example, vectors. WO95/07994, the disclosure of which is incorporated herein by reference.

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Where direct administration of the recombinant viral vector is contemplated, either in the form of naked nucleic acid or in the form of packaged particles carrying the nucleic acid this may be 20 done by any suitable means, for example oral administration or intravenous injection. From 105 to 108 c.f.u of virus represents a typical dose, which may be repeated for example weekly over a period of a few months. Administration of autologous or HLAmatched cells infected with the virus may be more convenient in This will generally be achieved by administering some cases. doses, for example from 105 to 106 cells per dose which may be repeated as described above.

The recombinant viral vector may further comprise nucleic acid capable of expressing an accessory molecule of the immune system designed to increase the immune response. Such a moleclue may be for example and interferon, particularly interferon gamma, an interleukin, for example IL-1 $\alpha$ , IL-1 $\beta$  or IL-2, or an HLA class I or II moleclue. This may be particularly desirable where the vector is intended for use in the treatment of humans or animals 35 already infected with a mycobacteria and it is desired to boost the immune response.

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## E. Antibodies.

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The invention also provides monoclonal or polyclonal antibodies to polypeptides of the invention or fragments thereof. The invention further provides a process for the production of monoclonal or polyclonal antibodies to polypeptides of the invention. Monoclonal antibodies may be prepared by conventional hybridoma technology using the polypeptides of the invention or peptide fragments thereof, as immunogens. Polyclonal antibodies may also be prepared by conventional means which comprise inoculating a host animal, for example a rat or a rabbit, with a polypeptide of the invention or peptide fragment thereof and recovering immune serum.

In order that such antibodies may be made, the invention also provides polypeptides of the invention or fragments thereof haptenised to another polypeptide for use as immunogens in animals or humans.

For the purposes of this invention, the term "antibody", unless specified to the contrary, includes fragments of whole antibodies which retain their binding activity for a polypeptide of the invention. Such fragments include Fv, F(ab') and F(ab')<sub>2</sub> fragments, as well as single chain antibodies. Furthermore, the antibodies and fragments thereof may be humanised antibodies, e.g. as described in EP-A-239400.

Antibodies may be used in methods of detecting polypeptides of the invention present in biological samples (where such samples include the human or animal body samples, and environmental samples, mentioned above) by a method which comprises:

- (a) providing an antibody of the invention;
- (b) incubating a biological sample with said antibody under conditions which allow for the formation of an antibody-antigen complex; and
- (c) determining whether antibody-antigen complex comprising said antibody is formed.

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Antibodies of the invention may be bound to a solid support for example an immunoassay well, microparticle, dipstick or biosensor and/or packaged into kits in a suitable container along with suitable reagents, controls, instructions and the like.

5 Antibodies of the invention may be used in the detection, diagnosis and prognosis of diseases as descirbed above in relation to polypeptides of the invention.

## F. Compositions.

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The present invention also provides compositions comprising a polynucleotide or polypeptide of the invention together with a carrier or diluent. Compositions of the invention also include compositions comprising a nucleic acid, particularly and expression vector, of the invention. Compositions further include those carrying a recombinant virus of the invention. Such compositions include pharmaceutical compositions in which case the carrier or diluent will be pharmaceutically acceptable.

Pharmaceutically acceptable carriers or diluents include those used in formulations suitable for inhalation as well as oral, parenteral (e.g. intramuscular or intravenous or transcutaneous) administration. The formulations may conveniently be presented in unit dosage form and may be prepared by any of the methods well known in the art of pharmacy. Such methods include the step of bringing into association the active ingredient with the carrier which constitutes one or more accessory ingredients. In general the formulations are prepared by uniformly and intimately bringing into association the active ingredient with liquid carriers or finely divided solid carriers or both, and then, if necessary, shaping the product.

For example, formulations suitable for parenteral administration include aqueous and non-aqueous sterile injection solutions which may contain anti-oxidants, buffers, bacteriostats and solutes which render the formulation isotonic with the blood of the intended recipient, and aqueous and non-aqueous sterile suspensions which may include suspending agents and thickening

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agents, and liposomes or other microparticulate systems which are designed to target the polynucleotide or the polypeptide of the invention to blood components or one or more organs, or to target cells such as M cells of the intestine after oral administration.

#### 5 G. Vaccines.

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In another aspect, the invention provides novel vaccines for the prevention and treatment of infections caused by Mptb, Mavs, other GS-containing pathogenic mycobacteria and Mtb in animals The term "vaccine" as used herein means an agent and humans. used to stimulate the immune system of a vertebrate, particularly a warm blooded vertebrate including humans, so as to provide protection against future harm by an organism to which the vaccine is directed or to assist in the eradication of an organism in the treatment of established infection. system will be stimulated by the production of cellular immunity desirably neutralizing antibodies, directed to epitopes found on or in a pathogenic mycobacterium which expresses any one of the ORFs of the invention. The antibody so produced may be any of the immunological classes, such as the immunoglobulins A, D, E, G or M. Vaccines which stimulate the production of IgA are interest since this is the principle immunoglobulin produced by the secretory system of warm-blooded animals, and the production of such antibodies will help prevent infection or colonization of the intestinal tract. However an IgM and IgG response will also be desirable for systemic infections such as Crohn's disease or tuberculosis.

Vaccines of the invention include polynucleotides of the invention or fragments thereof in suitable vectors and administered by injection of naked DNA using standard protocols. Polynucleotides of the invention or fragments thereof in suitable vectors for the expression of the polypeptides of the invention may be given by injection, inhalation or by mouth. Suitable vectors include M.bovis BCG, M.smegmatis or other mycobacteria, Corynebacteria, Salmonella or other agents according to established protocols.

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invention fragments Polypeptides of the orthereof substantially isolated form may be used as vaccines by injection, inhalation, oral administration or by transcutaneous application according to standard protocols. Adjuvants (such as Iscoms or polylactide-coglycolide encapsulation), cytokines such as IL-12 and other immunomodulators may be used for the selective enhancement of the cell mediated or humoral immunological responses. Vaccination with polynucleotides and/or polypeptides of the invention may be undertaken to increase the susceptibility of pathogenic mycobacteria to antimicrobial agents in vivo.

In instances wherein the polypeptide is correctly configured so as to provide the correct epitope, but is too small to be immunogenic, the polypeptide may be linked to a suitable carrier.

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A number of techniques for obtaining such linkage are known in the art, including the formation of disulfide linkages using Nsuccinimidyl-3-(2-pyridylthio) propionate (SPDP) and succinimidyl 4-(N-maleimido-methyl)cyclohexane-1-carboxylate (SMCC) obtained from Pierce Company, Rockford, Illinois, (if the peptide lacks a sulfhydryl group, this can be provided by addition of a cysteine residue). These reagents create a disulfide linkage between themselves and peptide cysteine residues on one protein and an amide linkage through the epsilon-amino on a lysine, or other free amino group in the other. A variety of such disulfide/amide-forming agents are known. See, for example, Immun Rev (1982) 62:185. Other bifunctional coupling agents form a thioether rather than a disulfide linkage. Many of these thioether-forming agents are commercially available and include reactive esters of 6-maleimidocaproic acid, 2-bromoacetic acid, 2-iodoacetic acid, 4-(N-maleimido-methyl)cyclohexane-1-carboxylic The carboxyl group can be activated by acid, and the like. combining them with succinimide or 1-hydroxyl-2-nitro-4-sulfonic acid, sodium salt. Additional methods of coupling antigens employs the rotavirus/"binding peptide" system described in EPO Pub. No. 259,149, the disclosure of which is incorporated herein by reference. The foregoing list is not meant to be exhaustive, and modifications of the named compounds can clearly be used.

Any carrier may be used which does not itself induce the production of antibodies harmful to the host. Suitable carriers are typically large, slowly metabolized macromolecules such as proteins; polysaccharides, such as latex functionalized Sepharose®, agarose, cellulose, cellulose beads and the like; polymeric amino acids, such as polyglutamic acid, polylysine, polylactide-coglycolide and the like; amino acid copolymers; and inactive virus particles. Especially useful protein substrates are serum albumins, keyhole limpet hemocyanin, immunoglobulin molecules, thyroglobulin, ovalbumin, tetanus toxoid, and other proteins well known to those skilled in the art.

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The immunogenicity of the epitopes may also be enhanced by preparing them in mammalian or yeast systems fused with or assembled with particle-forming proteins such as, for example, that associated with hepatitis B surface antigen. See, e.g., US-A-4,722,840. Constructs wherein the epitope is linked directly to the particle-forming protein coding sequences produce hybrids which are immunogenic with respect to the epitope. In addition, all of the vectors prepared include epitopes specific to HBV, having various degrees of immunogenicity, such as, for example, the pre-S peptide.

In addition, portions of the particle-forming protein coding sequence may be replaced with codons encoding an epitope of the invention. In this replacement, regions which are not required to mediate the aggregation of the units to form immunogenic particles in yeast or mammals can be deleted, thus eliminating additional HBV antigenic sites from competition with the epitope of the invention.

Vaccines may be prepared from one or more immunogenic polypeptides of the invention. These polypeptides may be expressed in various host cells (e.g., bacteria, yeast, insect, or mammalian cells), or alternatively may be isolated from viral preparations or made synthetically.

In addition to the above, it is also possible to prepare live vaccines of attenuated microorganisms which express one or more

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recombinant polypeptides of the invention. Suitable attenuated microorganisms are known in the art and include, for example, viruses (e.g., vaccinia virus), as well as bacteria.

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The preparation of vaccines which contain an immunogenic polypeptide(s) as active ingredients, is known to one skilled in Typically, such vaccines are prepared as injectables, or as suitably encapsulated oral preparations and either liquid solutions or suspensions; solid forms suitable for solution in, or suspension in, liquid prior to injestion or injection may also be prepared. The preparation may also be emulsified, or the protein encapsulated in liposomes. The active immunogenic often mixed with excipients ingredients are which pharmaceutically acceptable and compatible with the active ingredient. Suitable excipients are, for example, water, saline, dextrose, glycerol, ethanol, or the like and combinations thereof. In addition, if desired, the vaccine may contain minor amounts of auxiliary substances such as wetting or emulsifying agents, pH buffering agents, and/or adjuvants which enhance the effectiveness of the vaccine. Examples of adjuvants which may be effective include but are not limited to: aluminum hydroxide, N-acetyl-muramyl-L-threonyl-D-isoglutamine (thr-MDP), N-acetylnor-muramyl-L-alanyl-D-isoqlutamine (CGP 11637, referred to as nor-MDP), N-acetylmuramyl-L-alanyl-D-isoglutaminyl-L-alanine-2-(1'-2'-dipalmitoy1-sn-glycero-3-hydroxyphosphoryloxy) -ethylamine (CGP 19835A, referred to as MTP-PE), and RIBI, which contains 25 three components extracted from bacteria, monophosphoryl lipid A, trehalose dimycolate and cell wall skeleton (MPL+TDM+CWS) in a 2% squalene/Tween® 80 emulsion. The effectiveness of an adjuvant may be determined by measuring the amount of antibodies directed against an immunogenic polypeptide containing 30 antiqueic sequence resulting from administration of polypeptide in vaccines which are also comprised of the various adjuvants.

The vaccines are conventionally administered parenterally, by injection, for example, either subcutaneously or intramuscularly. Additional formulations which are suitable for other modes of administration include suppositories, oral formulations or as

enemas. For suppositories, traditional binders and carriers may include, for example, polyalkylene glycols or triglycerides; such suppositories may be formed from mixtures containing the active ingredient in the range of 0.5% to 10%, preferably 1% - 2%. Oral formulations include such normally employed excipients as, for example, pharmaceutical grades of mannitol, lactose, starch, magnesium stearate, sodium saccharine, cellulose, magnesium carbonate, and the like. These compositions take the form of solutions, suspensions, tablets, pills, capsules, sustained release formulations or powders and contain 10% - 95% of active ingredient, preferably 25% - 70%.

The proteins may be formulated into the vaccine as neutral or salt forms. Pharmaceutically acceptable salts include the acid addition salts (formed with free amino groups of the peptide) and which are formed with inorganic acids such as, for example, hydrochloric or phosphoric acids, or such organic acids such as acetic, oxalic, tartaric, maleic, and the like. Salts formed with the free carboxyl groups may also be derived from inorganic bases such as, for example, sodium, potassium, ammonium, calcium, or ferric hydroxides, and such organic bases as isopropylamine, trimethylamine, 2-ethylamino ethanol, histidine, procaine, and the like.

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The vaccines are administered in a manner compatible with the in such amount as will formulation, and prophylactically and/or therapeutically effective. The quantity to be administered, which is generally in the range of  $5\mu g$  to  $250\mu g$ , of antigen per dose, depends on the subject to be treated, capacity of the subject's immune system to synthesize antibodies, mode of administration and the degree of protection desired. Precise amounts of active ingredient required to be administered may depend on the judgement of the practitioner and may be peculiar to each subject.

The vaccine may be given in a single dose schedule, or preferably in a multiple dose schedule. A multiple dose schedule is one in which a primary course of vaccination may be with 1-10 separate doses, followed by other doses given at subsequent time intervals

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required to maintain and or reenforce the immune response, for example, at 1-4 months for a second dose, and if needed, a subsequent dose(s) after several months. The dosage regimen will also, at least in part, be determined by the need of the individual and be dependent upon the judgement of the practitioner.

In a further aspect of the invention, there is provided an attenuated vaccine comprising a normally pathogenic mycobacteria which harbours an attenuating mutation in any one of the genes encoding a polypeptide of the invention. The gene is selected from the group of ORFs A, B, C, D, E, F, G and H, including the homologous ORFs B, C, E and F in Mtb.

The mycobacteria may be used in the form of killed bacteria or as a live attenuated vaccine. There are advantages to a live attenuated vaccine. The whole live organism is used, rather than dead cells or selected cell components which may exhibit modified or denatured antigens. Protein antigens in the outer membrane will maintain their tertiary and quaternary structures. Therefore the potential to elicit a good protective long term immunity should be higher.

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The term "mutation" and the like refers to a genetic lesion in a gene which renders the gene non-functional. This may be at either the level of transcription or translation. The term thus envisages deletion of the entire gene or substantial portions thereof, and also point mutations in the coding sequence which result in truncated gene products unable to carry out the normal function of the gene.

A mutation introduced into a bacterium of the invention will generally be a non-reverting attenuating mutation. Non-reverting means that for practical purposes the probability of the mutated gene being restored to its normal function is small, for example less than 1 in 10<sup>6</sup> such as less than 1 in 10<sup>9</sup> or even less than 1 in 10<sup>12</sup>.

An attenuated mycobacteria of the invention may be in isolated form. This is usually desirable when the bacterium is to be used for the purposes of vaccination. The term "isolated" means that the bacterium is in a form in which it can be cultured, processed or otherwise used in a form in which it can be readily identified and in which it is substantially uncontaminated by other bacterial strains, for example non-attenuated parent strains or unrelated bacterial strains. The term "isolated bacterium" thus encompasses cultures of a bacterial mutant of the invention, for example in the form of colonies on a solid medium or in the form of a liquid culture, as well as frozen or dried preparations of the strains.

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In a preferred aspect, the attenuated mycobacterium further comprises at least one additional mutation. This may be a mutation in a gene responsible for the production of products essential to bacterial growth which are absent in a human or animal host. For example, mutations to the gene for aspartate semi-aldehyde dehydrogenase (asd) have been proposed for the production of attenuated strains of Salmonella. The asd gene is described further in Gene (1993) 129; 123-128. A lesion in the the enzyme aspartate  $\beta$ -semialdehyde encoding dehydrogenase would render the organism auxotrophic for the essential nutrient diaminopelic acid (DAP), which can be provided exogenously during bulk culture of the vaccine strain. this compound is an essential constituent of the cell wall for gram-negative and some gram-positive organisms and is absent from mammalian or other vertebrate tissues, mutants would undergo lysis after about three rounds of division in such tissues. Analogous mutations may be made to the attenuated mycobacteria of the invention.

In addition or in the alternative, the attenuated mycobacteria may carry a recA mutation. The recA mutation knocks out homologous recombination - the process which is exploited for the construction of the mutations. Once the recA mutation has been incorporated the strain will be unable to repair the constructed deletion mutations. Such a mutation will provide attenuated strains in which the possibility of homologous recombination to

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with DNA from wild-type strains has been minimized. RecA genes have been widely studied in the art and their sequences are available. Further modifications may be made for additional safety.

The invention further provides a process for preparing a vaccine composition comprising an attenuated bacterium according to the invention process comprises (a) inoculating a culture vessel containing a nutrient medium suitable for growth of said bacterium; (b) culturing said bacterium; (c) recovering said bacteria and (d) mixing said bacteria with a pharmaceutically acceptable diluent or carrier.

Attenuated bacterial strains according to the invention may be constructed using recombinant DNA methodology which is known per se. In general, bacterial genes may be mutated by a process of targeted homologous recombination in which a DNA construct containing a mutated form of the gene is introduced into a host bacterium which it is desired to attenuate. The construct will recombine with the wild-type gene carried by the host and thus the mutated gene may be incorporated into the host genome to provide a bacterium of the present invention which may then be isolated.

The mutated gene may be obtained by introducing deletions into the gene, e.g by digesting with a restriction enzyme which cuts the coding sequence twice to excise a portion of the gene and then religating under conditions in which the excised portion is not reintroduced into the cut gene. Alternatively frame shift mutations may be introduced by cutting with a restriction enzyme which leaves overhanging 5' and 3' termini, filling in and/or trimming back the overhangs, and religating. Similar mutations may be made by site directed mutagenesis. These are only examples of the types of techniques which will readily be at the disposal of those of skill in the art.

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Various assays are available to detect successful recombination. In the case of attenuations which mutate a target gene necessary for the production of an essential metabolite or catabolite

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compound, selection may be carried out by screening for bacteria unable to grow in the absence of such a compound. Bacteria may also be screened with antibodies or nucleic acids of the invention to determine the absence of production of a mutated gene product of the invention or to confirm that the genetic lesion introduced - e.g. a deletion - has been incorporated into the genome of the attenuated strain.

The concentration of the attenuated strain in the vaccine will be formulated to allow convenient unit dosage forms to be prepared. Concentrations of from about 10<sup>4</sup> to 10<sup>9</sup> bacteria per ml will generally be suitable, e.g. from about 10<sup>5</sup> to 10<sup>8</sup> such as about 10<sup>6</sup> per ml. Live attenuated organisms may be administered subcutaneously or intramuscularly at up to 10<sup>8</sup> organisms in one or more doses, e.g from around 10<sup>5</sup> to 10<sup>8</sup>, e.g about 10<sup>6</sup> or 10<sup>7</sup> organisms in a single dose.

The vaccines of the invention may be administered to recipients to treat established disease or in order to protect them against diseases caused by the corresponding wild type mycobacteria, such as inflammatory diseases such as Crohn's disease or sarcoidosis in humans or Johne's disease in animals. The vaccine may be administered by any suitable route. In general, subcutaneous or intramuscular injection is most convenient, but oral, intranasal and colorectal administration may also be used.

The following Examples illustrates aspects of the invention.

#### 25 EXAMPLE 1

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Tests for the presence of the GS identifier sequence were performed on  $5\mu$ l bacterial DNA extracts (25  $\mu$ g/ml to 500  $\mu$ g/ml) using polymerase chain reaction based on the oligonucleotide primers 5'-GATGCCGTGAGGAGGTAAAGCTGC-3' (Seq ID No. 40) and 5'-GATACGGCTCTTGAATCCTGCACG-3' (Seq ID No. 41) from within the identifier DNA sequences (Seq.ID Nos 1 and 2). PCR was performed for 40 cycles in the presence of 1.5 mM magnesium and an annealing temperature of  $58^{\circ}$ C. The presence or absence of the correct amplification product indicated the presence or absence

of GS identifier sequence in the corresponding bacterium. identifier sequence is shown to be present in all the laboratory and field strains of Mptb and Mavs tested. This includes Mptb isolates 0025 (bovine CVL Weybridge), 0021 (caprine, Moredun), (bovine, Moredun), 0139 (human, Chiodini 1984), 0209, 0022 0208, 0211, 0210, 0212, 0207, 0204, 0206 (bovine, Whipple 1990). All Mptb strains were IS900 positive. The Mavs strains include 0010 and 0012 (woodpigeon, Thorel) 0018 (armadillo, Portaels) and 0034, 0037, 0038, 0040 (AIDS, Hoffner). All Mavs strains were IS902 positive. One pathogenic M.avium strain 0033 (AIDS, Hoffner) also contained GS identifier sequence. GS identifier sequence is absent from other mycobacteria including other M.avium, M.malmoense, M.szulgai, M.gordonae, M. chelonei, M. fortuitum, M. phlei, as well as E. coli, S. areus, Nocardia sp, Streptococcus sp. Shigella sp. Pseudomonas sp.

## Example 2:

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To obtain the full sequence of GS in Mavs and Mptb we generated a genomic library of Mavs using the restriction endonuclease EcoRI and cloning into the vector pUC18. This achieved a representative library which was screened with 32P-labelled identifier sequence yielding a positive clone containing a 17kbp We constructed a restriction map of this insert and identified GS as fragments unique to Mavs and Mptb and not occurring in laboratory strains of M.avium. These fragments were sub-cloned into pUC18 and pGEM4Z. We identified GS contained within an 8kb region. The full nucleotide sequence was determined for GS on both DNA strands using primer walking and automated DNA sequencing. DNA sequence for GS in Mptb was obtained using overlapping PCR products generated using PwoDNA polymerase, a proofreading thermostable enzyme. The final DNA sequences were derived using the University of Wisconsin GCG qel assembly software package.

## Example 3:

The DNA sequence of GS in Mavs and Mptb was found to be more than 99% homologous. The ORFs encoded in GS were identified using GeneRunner and DNAStar computer programmes. Eight ORFs were identified and designated GSA, GSB, GSC, GSD, GSE, GSF, GSG

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Database comparisons were carried out against the GenEMBL Database release version 48.0 (9/96), using the BLAST and BLIXEM programmes. GSA and GSB encoded proteins of 13.5kDa and 30.7kDa respectively, both of unknown functions. GSC encoded a protein of 38.4kDa with a 65% homology to the amino acid sequence of rfbD of V.cholerae, a 62% amino acid sequence homology to gmd of E.coli and a 58% homology to gca of Ps.aeruginosa which are all GDP-D-mannose dehydratases. Equivalent gene products in H.influenzae, S.dysenteriae, Y.enterocolitica, N.gonorrhoea, K.pneumoniae and Salmonella enterica are all involved in 'O'-antigen processing known to be linked to pathogenicity. GSD encoded a protein of 37.1kDa which showed 58% homology at the DNA level to wcaG from a gene involved in the synthesis and regulation of capsular polysaccharides, also related to pathogenicity. was found to have a > 30% amino acid homology to rfbT of V. cholerae, involved in the transport of specific LPS components across the cell membrane. In V. cholerae the gene product causes a seroconversion from the Inaba to the Ogawa 'epidemic' strain. GSF encoded a protein of 30.2kDa which was homologous in the range 25-40% at the amino acid level to several glucosyl transferases such as rfpA of K.pneumoniae, rfbB of K.pneumoniae, lgtD of H.influenzae, 1si of N.gonorrhoae. In E.coli an equivalent gene galE adds  $\beta$ -1-3 N-acetylglucosamine to galactose, the latter only found in 'O' and 'M' antigens which are also related to pathogenicity. GSH comprising the ORFs GSH1 and GSH2 encodes a protein totalling about 60kDa which is a putative transposase with a 40 - 43% homology at the amino acid level to the equivalent gene product of IS21 in E.coli. This family of insertion sequences is broadly distributed amongst gram negative bacteria and is responsible for mobility and transposition of genetic elements. An IS21- like element in B. fragilis is split either side of the  $\beta$ -lactamase gene controlling its activation and expression. We programmed an E.coli S30 cell-free extract with plasmid DNA containing the ORF GSH under the control of a lac promoter in the presence of a 35S-methionine, demonstrated the translation of an abundant 60kDa protein. The proteins homologous to GS encoded in other organisms are in general highly antigenic. Thus the proteins encoded by the ORFs

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in GS may be used in immunoassays of antibody or cell mediated diagnosing infections immuno-reactivity for caused mycobacteria, particularly Mptb, Mavs and Mtb. Enhancement of host immune recognition of GS encoded proteins by vaccination using naked specific DNA or recombinant GS proteins, may be used in the prevention and treatment of infections caused by Mptb, Mavs and Mtb in humans and animals. Mutation or deletion of all or some of the ORFs A to H in GS may be used to generate attenuated strains of Mptb, Mavs or Mtb with lower pathogenicity for use as living or killed vaccines in humans and animals. Such vaccines are particularly relevant to Johne's disease in animals, to diseases caused by Mptb in humans such as Crohn's disease, and to the management of tuberculosis especially where the disease is caused by multiple drug-resistant organisms.

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### SEQUENCE LISTING

Seq. ID No.1

	5*- 1	GATCCAACTA	AACCCGATGG	AACCCCGCGC	AAACTATTGG	ACGTCTCCGC	GCTACGCAGT
	61	TGGGTTGGCG	CCCGCGAATC	GCACTGAAAG	AGGGCATCGA	TGCAACGGTG	TCGTGGTACC
5	121	GCACAAATGC	CGATGCCGTG	AGGAGGTAAA	GCTGCGGGCC	GGCCGATGTT	ATCCCTCCGG
	181	CCGGACGGGT	AGGGCGACCT	GCCATCGAGT	GGTACGGCAG	TCGCCTGGCC	GGCGAGGCGC
	241	ATGGCCTATG	TGAGTATCCC	ATAGCCTGGC	TTGGCTCGCC	CCTACGCATT	ATCAGTTGAC
	301	CGCTTTCGCG	CCACGTCGCA	GGCTTGCGGC	AGCATCCCGT	TCAGGTCTCC	TCATGGTCCG
	361	GTGTGGCACG	ACCACGCAAG	CTCGAACCGA	CTCGTTTCCC	AATTTCGCAT	GCTAATATCG
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•	481	GCGCCACTCC	GAACGAAAGC	GCCTATTAGT	AAACCAAGTC	GAAGCATACG	GAGTCAACGT
	541	TGTTATTGAT	GTCGGTGCTA	ACTCCGGCCA	GTTCGGTAGC	GCTTTGCGTC	GTGCAGGATT
	601	CAAGAGCCGT	ATCGTTTCCT	TTGAACCTCT	TTCGGGGCCA	TTTGCGCAAC	TAACGCGCAA
	661	GTCGGCATCG	GATC -3'				

15 Seq. ID No.2

661 GGTTTAGTTG GATC -3'

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5'-1 GATCGATGC CGACTTGCGC GTTAGTTGCG CAAATGGCCC CGAAAGAGGT TCAAAGGAAA
61 CGATACGGCT CTTGAATCCT GCACGACGCA AAGCGCTACC GAACTGGCCG GAGTTAGCAC
121 CGACATCAAT AACAACGTTG ACTCCGTATG CTTCGACTTG GTTTACTAAT AGGCGCTTTC
181 GTTCGGAGTG GCGCAGCATC TCGGTGCTAA CGTTACGAGC CATCAAGGCG GCGTTGCGCA
241 AAAAATCCAT CGAGCGATAT TAGCATGCGA AATTGGGAAA CGAGTCGGTT CGAGCTTGCG
301 TGGTCGTGCC ACACCGGACC ATGAGGAGAC CTGAACGGGA TGCTGCCGCA AGCCTGCGAC
361 GTGGCGCGAA AGCGGTCAAC TGATAATGCG TAGGGGCGAG CCAAGCCAGG CTATGGGATA
421 CTCACATAGG CCATGCGCCT CGCCGGCCAG GCGACTGCCG TACCACTCGA TGGCAGGTCG
481 CCCTACCCGT CCGGCCGGAG GGATAACATC GGCCGGCCCG CAGCTTTACC TCCTCACGGC
541 ATCGGCATT GTGCGGTACC ACGACACCGT TGCATCGATA GTTTGCGCGG GGTTCCATCG
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	201	CGGCGATGAA	AATGACGTCC	GCGTGCTCGA	TTCCGCGTTG	CCGGTCGGTG
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	801		CTGAACGATC			
	851		GCGCTATGGG			
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	951	-	TCTCAGCCCG			
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	1251		TCGCTGGTAG			
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	1351		TCGACCGCGT			
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	1551		TTGCGAGCGA			
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	2001					TATCGACCCG
	2051					TCAGCTTTGA
	2101					ATCCGACTTC
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	2251					TCGTACTGGA
	2301					GAATGGCATC
	2351					TGACCCGAAA
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	2451					GCCCGAATAT
	2501					ATGACTACGT
	2501	GI COMPROSE				

16147 = Start of SEQ 10 4 1 MAR.

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		2651	TATTTGCGTC	CCACCGAGGT	CGATTCGCTA	GTAGGAGATG	CCGACAAGGC
		2701	GGCCCAGTCA	CTCGGCTGGA	AAGCTTCGGT	TCATACTGGT	GAACTCGCGC
	5	2751	GCATCATGGT	GGACGCGGAC	ATCGCCGCGT	TGGAGTGCGA	TGGCACACCA
		2801	TGGATCGACA	CGCCGATGTT	GCCTGGTTGG	GGCAGAGTAA	GTTGACGACT
		2851	ACACCTGGGC	CTCTGGACCG	CGCAACGCCC	GTGTATATCG	CCGGTCATCG
		2901	GGGGCTGGTC	GGCTCAGCGC	TCGTACGTAG	ATTTGAGGCC	GAGGGGTTCA
		2951	CCAATCTCAT	TGTGCGATCA	CGCGATGAGA	TTGATCTGAC	GGACCGAGCC
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		3101	TCTTGTCCGA	AAACCTCCGA	ATCCAGACCA	ATTTGCTCGA	CGCAGCTGTC
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		4151	AAAGCGCCTA				
		4201	TTGATGTCGG		-		
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	5301	GTCTATGGCG	ATGTTGTGAT	GCGTTCGACG	AAAAGCCGGC	ATGCCGGACC
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	6351	CCCCATTGTT	CGAAGGGCCA	ATGCGAGGCG	ATGGCCAGGG	AGCGGCGCTC
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	7051	CGAGCGGGTC	TCGACGGTGG	GCAGATCGAG	CACGAGTGCG	TCGCCGGCGG
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	7251	ATTTCAGTCG	GGTGTTGCCG	ATCGCAGCAG	CACCGACGAG	GAACTGCTGC
	7301	GCTTCGGTTC	CCAATGCGCA	GAATCGTTTC	TCTGCTTGGG	TTTTCGGGCG
	7351	AGGACCACGC	GAGGGTGCGG	GTCTGGGTCC	GTCGTAGTGT	TCATCGAGGA
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	7551	GGATGCACGA	GAGGCCGTCG	ACCTTACGGC	GCACCGACCC	CGAGCCGATC
	7601	GTCGGCCGCA	GCGAGGGCAG	CTCCCTCAAG	ACGGTGCGCT	CGTCAACCAA
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	7751	TCACCGGCTA	ACGCAGCTTC	GGTCAGCAGC	GGCACCGCAA	GGTCGTCCTG
	7801	AGCGTAGCCA	CAGAGGTTCT	CCACGATGCC	CTTCGATTGC	GGATCCGCAC

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7951 CATCCGGTCG GCCAGGATCT TGGCCGGAAC CCCACCGATC GCCTC

Seq. ID No.4

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	661	GCGCGGCCAA	GGTCTTCTCG	TACTGGACGA	CTCGCAACTA	TCGAGAGGCG	TACGGATTAT
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	781	CCCGAAAGAT	CACGCGTGCC	GTGGCGCGCA	TCCGAGCTGG	CGTCCAATCG	GAGGTCTATA
	841	TGGGCAACCT	CGATGCGATC	CGCGACTGGG	GCTACGCGCC	CGAATATGTC	GAGGGGATGT
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	961	CCGTACGTGA	GTTCGCTCAA	GCTGCTTTTG	ACCACGTCGG	GCTCGACTGG	CAAAAGCACG
	1021	TCAAGTTTGA	CGACCGCTAT	TTGCGCCCCA	CCGAGGTCGA	TTCGCTAGTA	GGAGATGCCG
	1081	ACAGGGCGGC	CCAGTCACTC	GGCTGGAAAG	CTTCGGTTCA	TACTGGTGAA	CTCGCGCGCA
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	1321	TGAGGCCGAG	GGGTTCACCA	ATCTCATTGT	GCGATCACGC	GATGAGATTG	ATCTGACGGA
	1381	CCGAGCCGCA	ACGTTTGATT	TTGTGTCTGA	GACAAGACCA	CAGGTGATCA	TCGATGCGGC
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	1561	TTTCCTCGGT	TCGTCATGCA	TCTACCCGAA	GTACGCTCCG	CAACCTATCC	ACGAGAGTGC
	1621	TTTATTGACT	GGCCCTTTGG	AGCCCACCAA	${\tt CGACGCGTAT}$	GCGATCGCCA	AGATCGCCGG
	1681	TATCCTGCAA	GTTCAGGCGG	TTAGGCGCCA	ATATGGGCTG	GCGTGGATCT	CTGCGATGCC
	1741	GACTAACCTC	TACGGACCCG	GCGACAACTT	CTCCCCGTCC	GGGTCGCATC	TCTTGCCGGC
35	1801	GCTCATCCGT	CGATATGAGG	AAGCCAAAGC	TGGTGGTGCA	GAAGAGGTGA	CGAATTGGGG
	1861	GACCGGTACT	CCGCGGCGCG	AACTTCTGCA	TGTCGACGAT	CTGGCGAGCG	CATGCCTGTT
	1921	CCTTTTGGAA	CATTTCGATG	GTCCGAACCA	CGTCAACGTG	GGCACCGGCG	TCGATCACAG
	1981	CATTAGCGAG	ATCGCAGACA	TGGTCGCTAC	GGCGGTGGGC	TACATCGGCG	DITECACAAA
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40	2101	GTTGGGTTGG	CGCCCGCGAA	TCGCACTGAA	AGACGGCATC	GATGCAACGG	TGTCGTGGTA
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	2221	GGCCGGACGG	GTAGGGCGAC	CTGCCATCGA	GTGGTACGGC	AGTCGCCTGG	CCGGCGAGGC
	2281	GCATGGCCTA	TGGGAGTATC	CCATAGCCTG	GCTTGGCTCG	CCCCTACGCA	TTATCAGTTG
	2341	ACCGCTTTCG	CGCCAGCTCG	CAGGCTCGCG	GCAGCATCCC	GTTCAGGTCT	CCTCATGGTC
45	2401	CGGTGTGGCA	CGACCACGCA	AGCTCGAACC	GACTCGTTTC	CCAATTTCGC	ATGCTAATAT
	2461	CGCTCGATGG	ATTTTTTGCG	CAACGCCGGC	TTGATGGCTC	GTAACGTTAG	CACCGAGATG
	2521	CTGCGCCACT	TCGAACGAAA	GCGCCTATTA	GTAAACCAAT	TCAAAGCATA	CGGAGTCAAC
	2581	GTTGTTATTG	ATGTCGGTGC	TAACTCCGGC	CAGTTCGGTA	GCGCTTTGCG	TCGTGCAGGA
							ACTAACGCGC
50	2701	GAGTCGGCAT	CGGATCCACT	ATGGGAGTGT	CACCAGTATG	CCCTAGGCGA	CGCCGATGAG

PCT/GB96/03221

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Seq. ID No.7

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Seq. ID No.13

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Seq. ID No.17

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151 PTDVTFLKIDVQGFEKQVITGSKSTLNESC

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## 10 Seq. ID No.19

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### Seq. ID No.20

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91 Y A L G D A D E T I T I N V A G N A G A S S S V L P M L K S
121 H Q D A F P P A N Y I G T E D V A I H R L D S V A S E F L N
151 P T D V T F L K I D V Q G F E K Q V I A G S K S T L N E S C
181 V G M Q L E L S F I P L Y E G D M L I H E A L E L V Y S L G
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Seq. ID No.22

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121 G D V V M R S T K S R H A G P F D L D R L L F E T N L C H Q

151 S I F Y R R E L F D G I G P Y N L R Y R V W A D W D F N I R

181 C F S N P A L I T R Y M D V V I S E Y N D M T G F S M R Q G

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241 K E N R R L A L R T R L I R V K A V S K E R S A E P

Seq. ID No.23

1 atgactgogo cagtgttctc gataattatc cotaccttca atgcagoggt gacgctgcaa 61 gcctgcctcg gaagcatcgt cgggcagacc taccgggaag tggaagtggt ccttgtcgac 121 ggcggttcga ccgatcggac cctcgacatc gcgaacagtt tccgcccgga actcggctcg 30 181 cgactggteg ttcacagegg gecegatgat ggeecetaeg aegecatgaa cegeggegte 241 ggcgtageca caggcgaatg ggtacttttt ttaggcgccg acgacaccct ctacgaacca 301 accargitgg cocaggiage egetittete ggegaecatg eggeaageca tettgietat 361 qqcqatqttg tgatgcgttc gacgaaaagc cggcatgccg gacctttcga cetegaccgc 421 ctcctatttg agacgaattt gtgccaccaa tcgatctttt accgccgtga gcttttcgac 35 481 ggcatcggcc cttacaacct gcgctaccga gtctgggcgg actgggactt caatattegc 541 tgcttctcca acceggeget gattaccege tacatggaeg tegtgatttc egaatacaac 601 gacatgaccg gcttcagcat gaggcagggg actgatmaag agttcagama acggctgcca 661 atgtacttct gggttgcagg gtgggagact tgcaggcgca tgctggcgtt tttgaaagac 721 aaggagaato geegtetgge ettgegtaeg eggttgataa gggttaagge egteteeaaa 40 781 gaacgaageg cagaacegta g

5

10

Seq. ID No.24

1 MTAPVFSIIIPTFNAAVTLQACLGSIVGQT
31 YREVEVVLVDGGSTDRTLDIANSFRPELGS
61 RLVVHSGPDDGPYDAMNRGVGVATGEWVLF
91 LGADDTLYEPTTLAQVAAFLGDHAASHLVY
121 GDVVMRSTKSRHAGPFDLDRLLFETNLCHQ
151 SIFYRRELFDGIGPYNLRYRVWADWDFNIR
181 CFSNPALITRYMDVVISEYNDMTGFSMRQG
211 TDKEFRKRLPMYFWVAGWETCRRMLAFLKD

Seg. ID No.25

1 gtggccagca gaagtcccca ctccgctgcg ggtggttggc taattcttgg cggctccctt 61 cttgtggtcg gcgtggcgca tccggtagga ctcgccggag gtgacgacga tgctggcgtg 121 gtgcaqcage cgatcgagga tgctggcggc ggtggtgtgc tcgggcagga atcgccccca 15 181 ttgttegaag ggccaatgcg aggcgatggc cagggagcgg cgctcgtagc cggcagccac 241 gagccggaac aacagttgag tcccggtgtc gtcgagcggg gcgaagccga tctcgtccaa 301 gatgaccaga tccgcgcgga gcagggtgtc gatgatettg ccgacggtgt tgtcggccag 361 geogeggtag aggacetega teaggtegge ggeggtgaag tageggaett tgaateegge 421 gtggacggca gcgtgcccgc agccgatgag caggtgactt ttgcccgtac caggtgggcc 20 481 aatgacegee aggttetgtt gtgeeegaat ceatteeagg etegacaggt agtegaaegt 541 ggctgeggtg atcgacgatc eggtgacgtc gaaccegtcg agggtcttgg tgaccgggaa 601 ggetgeggee ttgagaeggt tggeggtgtt ggaggeateg egggeagega teteggeete 661 aaccaacgte egeaggatet eeteeggtgt eeagegttge gtettggega ettgeaacac 721 ctoggoggog ttgcggcgca ccgtggccag cttcaaccgc cgcagcgccg cgtcaaggtc 781 agcagccage ggtgccgccg aggacggtgc caccggcttg gcagcggtgg tcatgaggcc 25 841 gtcccgtcgg tggtgttgat cttgtag

Seq. ID No.26

	1	atgggctgcc	tcaaaggtgg	tgtcgtcgcc	aatgttgttg	ttccaacacc	ggattatgtg
	61	cgattcgcgt	cccactatgg	cttegtteeg	gacttctgcc	acggtgcgga	teegeaateg
	. 121	aagggcatcg	tggagaacct	ctgtggctac	gctcaggacg	accttgcggt	gccgctgctg
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	241	ctatggtgcg	ccgaggtcaa	tgccacggtc	cacteggaga	tetgegeegt	gcccaacgat
	301	cgcttggttg	acgagcgcac	cgtcttgagg	gagetgeect	cgctgcggcc	gacgatcggc
	361	tcggggtcgg	tgcgccgtaa	ggtcgacggc	ctctcgtgca	tccgttacgg	ctcagctcgt
	421	tactcggtgc	ctcagcggct	cgtcggtgcc	accgtggcgg	tggtggttga	tcatggcgcc
10	481	ctgatcctgt	tggaacctgc	gaccggtgtg	atcgtggccg	agcacgagct	cgtcagccca
	541	ggtgaggtgt	ccatcctcga	tgaacactac	gacggaccca	gacccgcacc	ctcgcgtggt
	601	cctcgcccga	aaacccaagc	agagaaacga	ttctgcgcat	tgggaaccga	agegeageag
	661	ttactagtag	gtgctgctgc	gatcggcaac	accegaetga	aatccgaact	cgacattctg
	721	ctcggccttg	gegeegeeca	cggcgaacag	gctttgattg	acgcgctgcg	ccgggcggtt
15	781	gcgtttcgcc	ggttccgcgc	tgccgacgtg	egetegatee	tggccgccgg	cgccggcacc
	841	ccacaacccc	acccaccaa	cgacgcactc	gtgctcgatc	tgcccaccgt	cgagacccgc
	901	tcgttggagg	cctacaagat	caacaccacc	gacgggacgg	cctcatgacc	accgctgcca
	961	agceggtgge	accgtcctcg	geggcacege	tggctgctga	ccttgacgcg	gegetgegge
	1021	ggttgaagct	ggccacggtg	cgccgcaacg	ccgccgaggt	gttgcaagtc	gccaagacgc
20	1081	aacgctggac	accggaggag	atcctgcgga	cgttggttga	ggccgagatc	gctgcccgcg
	1141	atgcctccaa	caccgccaac	cgtctcaagg	ccgcagcctt	cccggtcacc	aagaccctcg
	1201	acgggttcga	cgtcaccgga	tegtegatea	ccgcagccac	gttcgactac	ctgtcgagcc
	1261	tggaatggat	tegggeaeaa	cagaacctgg	cggtcattgg	cccacctggt	acgggcaaaa
	1321	gtcacctgct	catcggctgc	gggcacgctg	cegtecaege	cggattcaaa	gtccgctact
25	1381	tcaccgccgc	cgacctgatc	gaggtcctct	accgcggcct	ggccgacaac	accgtcggca
	1441	agatcatcga	caccctgctc	egegeggate	tggtcatctt	ggacgagatc	ggcttegccc
	1501	cgctcgacga	caccgggact	caactgttgt	teeggetegt	aacraccaac	tacgagegee
	1561	getecetgge	catcgcctcg	cattggccct	tcgaacaatg	ggggcgattc	ctgcccgagc
	1621	acaccaccgc	cgccagcatc	ctcgatcggc	tgctgcacca	egceageate	gtcgtcacct
30	1681	ccggcgagtc	ctaccggatg	egecaegeeg	accacaagaa	gggagccgcc	aagaattag

# Seq. ID No.28

1 M G C L K G G V V A N V V V P T P D Y V R F A S H Y G F V P 31 D F C H G A D P Q S K G I V E N L C G Y A Q D D L A V P L L 61 T E A A L A G E Q V D L R A L N A Q A Q L W C A E V N A T V 35 91 H S E I C A V P N D R L V D E R T V L R E L P S L R P T I G 121 S G S V R R K V D G L S C I R Y G S A R Y S V P Q R L V G A 151 T V A V V V D H G A L I L L E P A T G V I V A E H E L V S P 181 G E V S I L D E H Y D G P R P A P S R G P R P K T Q A E K R 211 F C A L G T E A Q Q F L V G A A A I G N T R L K S E L D I L A A G A G T P Q P R P A G D A L V L D L P T V E T R 301 S L E A Y K I N T T D G T A S

5

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Seq. ID No.29

1 M T T A A K P V A P S S A A P L A A D L D A A L R R L K L A
31 T V R R N A A E V L Q V A K T Q R W T P E E I L R T L V E A
61 E I A A R D A S N T A N R L K A A A F P V T K T L D G F D V
91 T G S S I T A A T F D Y L S S L E W I R A Q Q N L A V I G P
121 P G T G K S H L L I G C G H A A V H A G F K V R Y F T A A D
151 L I E V L Y R G L A D N T V G K I I D T L L R A D L V I L D
181 E I G F A F L D D T G T Q L L F R L V A A G Y E R R S L A I
211 A S H W P F E Q W G R F L P E H T T A A S I L D R L L H H A
241 S I V V T S G E S Y R M R H A D H K K G A A K N

Seq. ID No.30

1 gtgacgtctg ctccgaccgt ctcggtgata acgatctcgt tcaacgacct cgacgggttq 61 cagegcacgg tgaaaagtgt gegggegeaa egetaceggg gaegcatega geacategta 121 alogacggtg gcagcggcga cgacgtggtg gcatacctgt ccgggtgtga accaggcttc 15 181 gegtattggc agtccgagec cgacggcggg cggtacgacg cgatgaacca gggcategeg 241 caegeategg gtgatetgtt gtggttettg caeteegeeg ategttttte egggeeegae 301 gtggtageec aggeegtgga ggegetatec ggeaagggae eggtgteega attgtgggge 361 ttcgggatgg atcgtctcgt cgggctcgat cgggtgcgcg gcccgatacc tttcagcctg 421 ogcaaattoo tggooggcaa goaggttgtt cogcatoaag catcgttott oggatoatog 20 481 ctggtggcca agatcggtgg ctacgacctt gatttcggga tcgccgccga ccaggaattc 541 atattgoggg cogogotggt atgogagoog gtoacgatto ggtgtgtget gtgogagtto 601 gacaccacgg gcgtcggctc gcaccgggaa ccaagcgcgg tottcggtga totgcgccgc 661 atgggegaec ttcategeeg ctaccegttc gggggaagge gaatatcaca tgcctaccta 721 cgoggccggg agttctacgc ctacaacagt cgattctggg aaaacgtctt cacgcgaatg 25 781 tegaaatag

Seq. ID No.31

1 M T S A P T V S V I T I S F N D L D G L Q R T V K S V R A Q
31 R Y R G R I E H I V I D G G S G D D V V A Y L S G C E P G F
61 A Y W Q S E P D G G R Y D A M N Q G I A H A S G D L L W F L
30 91 H S A D R F S G P D V V A Q A V E A L S G K G P V S E L W G
121 F G M D R L V G L D R V R G P I P F S L R K F L A G K Q V V
151 P H Q A S F F G S S L V A K I G G Y D L D F G I A A D Q E F
181 I L R A A L V C E P V T I R C V L C E F D T T G V G S H R E
211 P S A V F G D L R R M G D L H R R Y P F G G R R I S H A Y L
35 241 R G R E F Y A Y N S R F W E N V F T R M S K

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1 gtgaagegag egeteateae eggaateace ggecaggaeg getegtatet egeogaactg
               61 ctgctggcca aggggtatga ggttcacggg ctcatccggc gcgcttcgac gttcaacacc
              121 tegeggateg ateaceteta egtegaceeg caccaacegg gegegegget gittetgeac
5
              181 tatggtgacc tgatcgacgg aacccggttg gtgaccctgc tgagcaccat cgaacccgac
              241 gaggtgtaca acctggcggc gcagtcacac gtgcgggtga gcttcgacga accegtgcac
              301 acceptaca ccacceptat gegatecate ceactetes aagcepttee ectetes
              361 gtgcactgcc gettetatea ggogtecteg teggagatgt teggegeete geogecaceg
              421 cagaacgage tgacgccgtt ctacccgcgg tcaccgtatg gcgccgccaa ggtctattcg
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              481 tactgggcga cccgcaatta tegcgaagcg tacggattgt tegccgttaa eggcatettg
              541 ttcaatcacg aatcaccgcg gcgcggtgag acgttcgtga cccgaaagat caccagggcc
              601 gtggcaegca tcaaggecgg tatccagtee gaggtctata tgggcaatet ggatgeggte
              661 cgcgactggg ggtacgcgcc cgaatacgtc gaaggcatgt ggcggatgct gcagaccgac
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              781 geogegtteg ageatgeogg tittggactgg cageagtacq tgaaattega ccaacgetat
              841 etgeggecca cegaggtgga ttegetgate ggegaegega ecaaggetge egaattgetg
              901 ggctggaggg cttcggtgca cactgacgag ttggctcgga tcatggtcga cgcggacatg
              961 geggegeteg agtgegaagg caageegteg ategacaage egatgatege eggeeggaca
             1021 tga
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20 Seq. ID No.33

1 M K R A L I T G I T G Q D G S Y L A E L L L A K G Y E V H G
31 L I R R A S T F N T S R I D H L Y V D P H Q P G A R L F L H
61 Y G D L I D G T R L V T L L S T I E P D E V Y N L A A Q S H
91 V R V S F D E P V H T G D T T G M G S M R L L E A V R L S R
121 V H C R F Y Q A S S S E M F G A S P P P Q N E L T P F Y P R
151 S P Y G A A K V Y S Y W A T R N Y R E A Y G L F A V N G I L
181 F N H E S P R R G E T F V T R K I T R A V A R I K A G I Q S
211 E V Y M G N L D A V R D W G Y A P E Y V E G M W R M L Q T D
241 E P D D F V L A T G R G F T V R E F A R A A F E H A G L D W
30 271 Q Q Y V K F D Q R Y L R P T E V D S L I G D A T K A A E L L
301 G W R A S V H T D E L A R I M V D A D M A A L E C E G K P W

Seq. ID No.34

1 atgaggetgg cocgtegege teggaacate ttgegtegea aeggeatega ggtgtegege 35 61 tactttgccg aactggactg ggaacgcaat ttcttgcgcc aactgcaatc gcatcgggtc 121 agtgccgtgc tcgatgtcgg ggccaattcg gggcagtacg ccaggggtct gcgcgcgcg 181 ggcttcgcgg gccgcatcgt ctcgttcgag ccgctgcccg ggccctttgc cgtcttgcag 241 egeagegeet ceaeggaeee gttgtgggaa tgeeggeget gtgegetggg egatgtegat 301 ggaaccatct cgatcaacgt cgccggcaac gagggcgcca gcagttccgt cttgccgatg 40 361 ttgaaacgac atcaggacgc ctttccacca gccaactacg tgggcgccca acgggtgccg 421 atacategae tegatteegt ggetgeagae gttetgegge ecaaegatat tgegttettg 481 aagategacg ttcaaggatt cgagaagcag gtgategegg gtggcgattc aacggtgcac 541 gaccgatgcg tcggcatgca gctcgagctg tctttccagc cgttgtacga gggtggcatg 601 ctcatecgcg aggegetega tetegtggat tegttggget ttaegetete gggattgeaa 45 661 cooggettea cogacceecg caacggeega atgetgeagg cogatggeat cttetteegg 721 ggcagcgatt ga

5

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1 M R L A R R A R N I L R R N G I E V S R Y F A E L D W E R N 31 F L R Q L Q S H R V S A V L D V G A N S G Q Y A R G L R G A 61 G F A G R I V S F E P L P G P F A V L Q R S A S T D P L W E 91 C R R C A L G D V D G T I S I N V A G N E G A S S S V L P M 121 L K R H Q D A F P P A N Y V G A Q R V P I H R L D S V A A D 151 V L R P N D I A F L K I D V Q G F E K Q V I A G G D S T V H 181 D R C V G M Q L E L S F Q P L Y E G G M L I R E A L D L V D 211 S L G F T L S G L Q P G F T D P R N G R M L Q A D G I F F R 241 G S D

Seq. ID No.36

1 gtgaaatcgt tgaaactcgc tegttteate gegegtageg cegeettega ggtttegege 61 cgetattetg agegagacet gaageaceag tttgtgaage aacteaaate gegeggggta 121 gatgtegttt tegatgtegg egecaactea ggacaataeg cegeeggeet cegeeggea 121 geatataagg geegeattgt etegttegaa eegetateeg gacegtttae gatettggaa 241 ageaaagegt caacggatee actttgggat tgeeggeage atgegttgg egattetgat 301 ggaaeggtta egateaatat egeaggaaae geeggteaga geagtteegt ettgeeattg 361 etgaaaagee ttgatteege ggegeeagaa tteteaggaa tegggtage egattetete 20 421 atacategae ttgatteegt ggegeeagaa tteteaggea tgaaeggtgt egetttete 20 481 aaggtegaeg teeaagget tgaaaageag gtgeteegee ggggeaaaate aaceatagat 541 gaceattgeg teggeatgea acteggaaeg teetteete 661 eetatteetg aageeetega teetgtgtat teettegget teaeggtae ggggtatgetg eggattgetg 661 eetatteetg tegatgeaa taatggtega atgttgeag eegatggeat etttteege 721 gaggaegatt gg

25 Seq. ID No.37

1 M K S L K L A R F I A R S A A F E V S R R Y S E R D L K H Q
31 F V K Q L K S R R V D V V F D F T V G A N S G Q Y A A G L R
61 R A A Y K G R I V S F E P L S G P F T I L E S K A S T D P L
91 W D C R Q H A L G D S D G T V T I N I A G N A G Q S S S V L
121 P M L K S H Q N A F P P A N Y V G T Q E A S I H R L D S V A
151 P E F L G M N G V A F L K V D V Q G F E K Q V L A G G K S T
181 I D D H C V G M Q L E L S F L P L Y E G G M L I P E A L D L
211 V Y S L G F T L T G L L P C F I D A N N G R M L Q A D G I F

1 atggtgcaga cgaaacgata cgccggcttg accgcagcta acacaaagaa agtcgccatg 61 geograceaa tgttttegat cateateece acettgaacg tggetgeggt attgeetgee 121 tgeetegaca geategeeeg teagacetge ggtgaetteg agetggtaet qgtegacqqe 5 181 ggctegaegg acgaaaccet cgacategec aacatttteg eccecaacet eggegagegg 241 ttgatcattc atcgcgacac cgaccagggc gtotacgacg ccatgaaccg cggcgtggac 301 otggocaccy gaacgtggtt getetttetg ggegeggaeg acagectgta cgaggetgae 361 accorgage gggrageege etteattage gaacacgage coagegatet ggtatatage 421 gacgtgatca tgcgctcaac caatttecgc tggggtggcg ccttcgacct cgaccgtctq 10 481 ttgttcaage gcaacatetg coatcaggeg atettetace gccgcggact ctteggeace 541 ateggteest assacsteeg stacegggte etggeegast gggastteaa tattegetge 601 ttttccaacc cagogetegt caccegetae atgeaegtgg tegttgcaag etacaacqaa 661 ttoggoggge tcageaatac gatogtogac aaggagtttt tgaagegget gcegatgtoc 721 acgagactog gcataagget ggtcatagtt ctggtgcgca ggtggccaaa ggtgatcage 15 781 agggccatgg taatgcgcac cgtcatttct tggcggcgcc gacgttag

Seq. ID No.39

1 M V Q T K R Y A G L T A A N T K K V A M A A P M F S I I I P
31 T L N V A A V L P A C L D S I A R Q T C G D F E L V L V D G
61 G S T D E T L D I A N I F A P N L G E R L I I H R D T D Q G
91 V Y D A M N R G V D L A T G T W L L F L G A D D S L Y E A D
121 T L A R V A A F I G E H E P S D L V Y G D V I M R S T N F R
151 W G G A F D L D R L L F K R N I C H Q A I F Y R R G L F G T
181 I G P Y N L R Y R V L A D W D F N I R C F S N P A L V T R Y
211 M H V V V A S Y N E F G G L S N T I V D K E F L K R L P M S
25 241 T R L G I R L V I V L V R R W P K V I S R A M V M R T V I S
271 W R R R R

Seq 40:

GATGCCGTGAGGAGGTAAAGCTGC

Seq 41:

30 GATACGGCTCTTGAATCCTGCACG

### CLAIMS

- 1. A polypeptide in substantially isolated form which comprises any one of the sequences selected from the group consisting of Seq.ID.No: 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 29, 31, 33, 35, 37 and 39, or a polypeptide substantially homologous thereto.
- 2. A polypeptide in substantially isolated form which comprises any one of the sequences selected from the group consisting of Seq.ID.No: 6, 8, 10, 12, 14, 16, 18, 20, 22, 24, 26, 28, 29, 31, 33, 35, 37 and 39.
- 3. A polypeptide which comprises a fragment of a polypeptide defined in claim 1 or 2, said fragment comprising at least 12 amino acids and an epitope.
- 4. A polynucleotide in substantially isolated form which encodes a polypeptide according to any one of claims 1 to 3.
- 5. A polynucleotide in substantially isolated form which is capable of selectively hybridizing to SEQ ID NO: 3 or 4 or a fragment thereof.
- 6. A polynucleotide fragment according to claim 5 which comprises a sequence selected from the group consisting of Seq.ID.No: 5, 7, 9, 11, 13, 15, 17, 19, 21, 23, 25 and 27, or a polynucleotide at least 90% homologous thereto.
- 7. A polynucleotide in substantially isolated form comprising a sequence selected from the group consisting of Seq.ID.No: 5, 7, 9, 11, 13, 15, 17, 19, 21, 23, 25 and 27.
- 8. A polynucleotide in substantially isolated form consiting essentially of a sequence selected from the group Seq ID Nos. 30, 32, 34, 36 and 38, or a polynucleotide at least 90% homologous thereto.

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- 9. A polynucleotide in substantially isolated form consiting essentially of a sequence selected from the group Seq ID Nos. 30, 32, 34, 36 and 38.
- 10. A polynucleotide probe which comprises a fragment of at least 15 nucleotides of a polynucleotide as defined in any one of claims 4 to 8, optionally carrying a revealing label.
- 11. A recombinant vector carrying a polynucleotide as defined in any one of claims 4 to 8.
- 12. An antibody capable of binding a polypeptide or fragment thereof as defined in any one of claims 1 to 3.
- 13. A test kit for detecting the presence or absence of a pathogenic mycobacterium in a sample which comprises a polynucleotide according to any one of claims 4 to 10, a polypeptide according to any one of claims 1 to 3, or an antibody according to claim 12.
- 14. A method of detecting the presence or absence of antibodies in an animal or human, against a pathogenic mycobacteria in a sample which comprises:
  - (a) providing a polypeptide according to any one of claims 1 to 3 comprising an epitope;
  - (b) incubating a biological sample with said polypeptide under conditions which allow for the formation of an antibody-antigen complex; and
  - (c) determining whether antibody-antigen complex comprising said polypeptide is formed.
- 15. A method of detecting the presence or absence of a polypeptide according to any one of claims 1 to 3 in a biological sample which method which comprises:
  - (a) providing an antibody according to claim 11;
  - (b) incubating a biological sample with said antibody under conditions which allow for the formation of an antibody-antigen complex; and

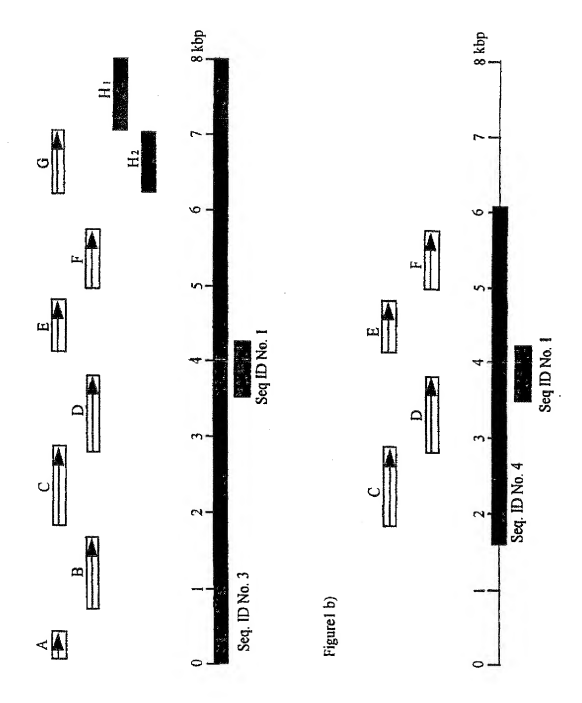
- 59 -

- (c) determining whether antibody-antigen complex comprising said antibody is formed.
- 16. A method of detecting the presence or absence of cell mediated immune reactivity in an animal or human, to a polypeptide according to claims 1 to 3 which method comprises
  - (a) providing a polypeptide according to any one of claims 1 to 3 comprising an epitope;
  - (b) incubating a cell sample with said polypeptide under conditions which allow for a cellular immune response such as release of cytokines or other mediator or reaction to occur; and
  - (c) detecting the presence of said cytokine or mediator or cellular response in the incubate.
- 17. A pharmaceutical composition comprising a polypeptide according to any one of claims 1 to 3 in a suitable carrier or diluent.
- 18. A composition according to claim 17 for use in the treatment or prevention of diseases caused by mycobacteria.
- 19. A method of treating or preventing mycobacterial disease in an animal or human caused by mycobacteria which express a polypeptide according to claims 1 to 3, which method comprises vaccinating or treating an animal or human with an effective amount of said polypeptide.
- 20. A method of treating or preventing mycobacterial diseases in animals or humans caused by mycobacteria containing the polynucleotide of SEQ ID NO: 3 or 4, which method comprises vaccinating or treating an animal or human with an effective amount of a polynucleotide according to claims 4 to 9, or a vector according to claim 11.
- 21. A method according to claims 19 or 20 for increasing the in vivo susceptibility of mycobacteria to antimicrobial drugs.

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- 22. A vaccine comprising a normally pathogenic mycobacteria, which pathogenicity is mediated in all or in part by the presence of the expression of a polypeptide as defined in any one of claims 1 to 3, which mycobacteria harbours an attenuating mutation in any one of said genes.
- 23. A vaccine according to claim 22 wherein the mycobacteria is selected from Mavs, Mptb and Mtb.

Figure 1 a)



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